

PATENT COOPERATION TREATY

PCT

INTERNATIONAL SEARCH REPORT

(PCT Article 18 and Rules 43 and 44)

Applicant's or agent's file reference BLP:106 PCT	FOR FURTHER ACTION	see Notification of Transmittal of International Search Report (Form PCT/ISA/220) as well as, where applicable, item 5 below.
International application No. PCT/US99/20776	International filing date (day/month/year) 10 September 1999 (10.09.1999)	(Earliest) Priority Date (day/month/year) 11 September 1998 (11.09.1998)
Applicant BERKSHIRE LABORATORIES, INC.		

This international search report has been prepared by this International Searching Authority and is transmitted to the applicant according to Article 18. A copy is being transmitted to the International Bureau.

This international search report consists of a total of 4 sheets.



It is also accompanied by a copy of each prior art document cited in this report.

1. Basis of the Report

a. With regard to the language, the international search was carried out on the basis of the international application in the language in which it was filed, unless otherwise indicated under this item.



the international search was carried out on the basis of a translation of the international application furnished to this Authority (Rule 23.1(b)).

b. With regard to any nucleotide and/or amino acid sequence disclosed in the international application, the international search was carried out on the basis of the sequence listing:



contained in the international application in written form.



filed together with the international application in computer readable form.



furnished subsequently to this Authority in written form.



furnished subsequently to this Authority in computer readable form.



the statement that the subsequently furnished written sequence listing does not go beyond the disclosure in the international application as filed has been furnished.



the statement that the information recorded in computer readable form is identical to the written sequence listing has been furnished.

2. ☐ Certain claims were found unsearchable (See Box I).

3. ☐ Unity of invention is lacking (See Box II).

4. With regard to the title,



the text is approved as submitted by the applicant.



the text has been established by this Authority to read as follows:

Please See Continuation Sheet

5. With regard to the abstract,



the text is approved as submitted by the applicant.



the text has been established, according to Rule 38.2(b), by this Authority as it appears in Box III. The applicant may, within one month from the date of mailing of this international search report, submit comments to this Authority.

6. The figure of the drawings to be published with the abstract is Figure No. 14



as suggested by the applicant.



None of the figures



because the applicant failed to suggest a figure.



because this figure better characterize the invention.

INTERNATIONAL SEARCH REPORT

International application No.

PCT/US99/20776

A. CLASSIFICATION OF SUBJECT MATTER

IPC(7) : A61B 17/22, 6/00; A61N 7/00

US CL : 600/427, 429, 439; 601/2, 3, 4; 73/579

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

U.S. : 600/427, 429, 439; 601/2, 3, 4; 607/97, 51; 604/22; 73/579

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)
EAST: resonant, resonance, biologic, biological, cell, acoustic, ultrasonic, ultrasound

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	US 3,774,717 A (CHODOROW) 27 November 1973 (27.11.1973) see abstract, Figures, column 1 line 35 - column 3 line 6	1-2, 23-25, 28, 34
---		10, 16-18, 21-22, 35
Y		
---		3-4, 19-20, 27, 36-39
A		1, 5-7, 9-11, 13
X	US 4,315,514 A (DREWES ET AL.) 16 February 1982 (16.02.1982), see abstract, Figures, column 2 lines 30-64.	8, 12, 14-18, 21-22, 35

Y		35
Y	US 5,413,550 A (CASTEL) 09 May 1995 (09.05.1995) see abstract, Figures, column 1 lines 5-16, column 6 line 55 - column 7 line 27	8, 10, 12, 14-15
Y	US 5,595,178 A (VOSS ET AL.) 21 January 1997 (21.01.1997) see abstract, Figures, column 1 lines 5-18, column 2 lines 5-29, column 3 line 47 - column 4 line 13, column 4 lines 29-44	25-26, 28-29, 32-34
X	US 5,777,228 A (TSUBOI ET AL.) 07 July 1998 (07.07.1998) see abstract, Figures, column 3 line 33 - column 9 line 20	

☒ Further documents are listed in the continuation of Box C.



See patent family annex.

* Special categories of cited documents:

"A" document defining the general state of the art which is not considered to be of particular relevance

"E" earlier application or patent published on or after the international filing date

"L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)

"O" document referring to an oral disclosure, use, exhibition or other means

"P" document published prior to the international filing date but later than the priority date claimed

"T"

later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention

"X"

document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone

"Y"

document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art

"&"

document member of the same patent family

Date of the actual completion of the international search

04 March 2000 (04.03.2000)

Date of mailing of the international search report

Name and mailing address of the ISA/US

Commissioner of Patents and Trademarks
Box PCT
Washington, D.C. 20231

Facsimile No. (703)305-3230

Authorized officer

Hezron Williams

Telephone No. 703-308-0956

INTERNATIONAL SEARCH REPORT

International application No.

PCT/US99/20776

C (Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X, P	US 5,886,263 A (NATH ET AL.) 23 March 1999 (23.03.1999) see abstract, Figures, column 1 line 33	25-26, 28-29, 32-34
---	- column 2 line 17	-----
A, P		27, 30-31, 36-38

INTERNATIONAL SEARCH REPORT

International application No.

PCT/US99/20776

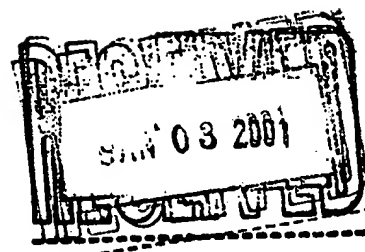
Continuation of Item 4 of the first sheet: METHODS FOR USING RESONANT ACOUSTIC ENERGY TO DETECT OR EFFECT STRUCTURES

PATENT COOPERATION TREATY

PCT

INTERNATIONAL PRELIMINARY EXAMINATION REPORT

(PCT Article 36 and Rule 70)



Applicant's or agent's file reference BLP:106 PCT		FOR FURTHER ACTION See Notification of Transmittal of International Preliminary Examination Report (Form PCT/IPEA/416)	
International application No. PCT/US99/20776	International filing date (day/month/year) 10 September 1999 (10.09.1999)	Priority date (day/month/year) 11 September 1998 (11.09.1998)	
International Patent Classification (IPC) or national classification and IPC IPC(7): A61B 17/22, 6/00; A61N 7/00 and US Cl.: 600/427, 429, 439; 601/2, 3, 4; 73/579			
Applicant BERKSHIRE LABORATORIES, INC.			
<p>1. This international preliminary examination report has been prepared by this International Preliminary Examining Authority and is transmitted to the applicant according to Article 36.</p> <p>2. This REPORT consists of a total of <u>6</u> sheets, including this cover sheet.</p> <p><input checked="" type="checkbox"/> This report is also accompanied by ANNEXES, i.e., sheets of the description, claims and/or drawings which have been amended and are the basis for this report and/or sheets containing rectifications made before this Authority (see Rule 70.16 and Section 607 of the Administrative Instructions under the PCT).</p> <p>These annexes consist of a total of <u>26</u> sheets.</p>			
<p>3. This report contains indications relating to the following items:</p> <p>I <input checked="" type="checkbox"/> Basis of the report</p> <p>II <input type="checkbox"/> Priority</p> <p>III <input type="checkbox"/> Non-establishment of report with regard to novelty, inventive step and industrial applicability</p> <p>IV <input type="checkbox"/> Lack of unity of invention</p> <p>V <input checked="" type="checkbox"/> Reasoned statement under Article 35(2) with regard to novelty, inventive step or industrial applicability; citations and explanations supporting such statement</p> <p>VI <input type="checkbox"/> Certain documents cited</p> <p>VII <input type="checkbox"/> Certain defects in the international application</p> <p>VIII <input type="checkbox"/> Certain observations on the international application</p>			
Date of submission of the demand 10 April 2000 (10.04.2000)		Date of completion of this report 11 December 2000 (11.12.2000)	
Name and mailing address of the IPEA/US Commissioner of Patents and Trademarks Box PCT Washington, D.C. 20231 Facsimile No. (703) 305-3230		Authorized officer Hezron Williams Telephone No. (703) 308-0956 <i>Pence</i>	

INTERNATIONAL PRELIMINARY EXAMINATION REPORT

International application No.

PCT/US99/20776

I. Basis of the report

1. With regard to the elements of the international application:*

- ☐ the international application as originally filed.
- ☒ the description:
pages 2, 14-21, 23-25, 27-37, 39-40, 42-46, 49-53, 55-62, 64-66 as originally filed
pages NONE, filed with the demand
pages 1, 3-13, 22, 26, 38, 41, 47-48, 54 and 63, filed with the letter of 22 September 2000 (22.09.2000)
- ☒ the claims:
pages NONE, as originally filed
pages NONE, as amended (together with any statement) under Article 19
pages NONE, filed with the demand
pages 67-73, filed with the letter of 22 September 2000 (22.09.2000)
- ☒ the drawings:
pages 1-28, as originally filed
pages NONE, filed with the demand
pages NONE, filed with the letter of _____
- ☒ the sequence listing part of the description:
pages NONE, as originally filed
pages NONE, filed with the demand
pages NONE, filed with the letter of _____

2. With regard to the language, all the elements marked above were available or furnished to this Authority in the language in which the international application was filed, unless otherwise indicated under this item.

These elements were available or furnished to this Authority in the following language _____ which is:

- ☐ the language of a translation furnished for the purposes of international search (under Rule 23.1(b)).
- ☐ the language of publication of the international application (under Rule 48.3(b)).
- ☐ the language of the translation furnished for the purposes of international preliminary examination (under Rules 55.2 and/or 55.3).

3. With regard to any nucleotide and/or amino acid sequence disclosed in the international application, the international preliminary examination was carried out on the basis of the sequence listing:

- ☐ contained in the international application in printed form.
- ☐ filed together with the international application in computer readable form.
- ☐ furnished subsequently to this Authority in written form.
- ☐ furnished subsequently to this Authority in computer readable form.
- ☐ The statement that the subsequently furnished written sequence listing does not go beyond the disclosure in the international application as filed has been furnished.
- ☐ The statement that the information recorded in computer readable form is identical to the written sequence listing has been furnished.

4. ☒ The amendments have resulted in the cancellation of:

- ☒ the description, pages NONE
- ☒ the claims, Nos. NONE
- ☒ the drawings, sheets/fig NONE

5. ☐ This report has been established as if (some of) the amendments had not been made, since they have been considered to go beyond the disclosure as filed, as indicated in the Supplemental Box (Rule 70.2(c)).**

* Replacement sheets which have been furnished to the receiving Office in response to an invitation under Article 14 are referred to in this report as "originally filed" and are not annexed to this report since they do not contain amendments (Rules 70.16 and 70.17).

** Any replacement sheet containing such amendments must be referred to under item 1 and annexed to this report.

INTERNATIONAL PRELIMINARY EXAMINATION REPORT

International application No.

PCT/US99/20776

V. Reasoned statement under Article 35(2) with regard to novelty, inventive step or industrial applicability; citations and explanations supporting such statement

1. STATEMENT

Novelty (N)	Claims <u>1-28, 30-31, 35-44</u>	YES
	Claims <u>29, 32-34</u>	NO
Inventive Step (IS)	Claims <u>1-28, 30-31, 35-44</u>	YES
	Claims <u>29, 32-34</u>	NO
Industrial Applicability (IA)	Claims <u>1-44</u>	YES
	Claims <u>NONE</u>	NO

2. CITATIONS AND EXPLANATIONS (Rule 70.7)

Please See Continuation Sheet

INTERNATIONAL PRELIMINARY EXAMINATION REPORT

International application No.
PCT/US99/20776

Supplemental Box

(To be used when the space in any of the preceding boxes is not sufficient)

V. 2. Citations and Explanations:

Claims 1-2 and 4-5 meet the criteria set out in PCT Article 33(2)-(4), because the prior art does not teach or fairly suggest a method for targeting a biologic structure to augment its function which comprises irradiating the biologic structure with acoustic energy to induce acoustic resonance therein.

Claims 3 and 6-7 meet the criteria set out in PCT Article 33(2)-(4), because the prior art does not teach or fairly suggest a method for targeting a biologic structure to affect at least one function which comprises irradiating the biologic structure with acoustic energy having a frequency near or at the resonant frequency of the biologic structure to induce acoustic resonance therein, and determining at least one acoustic signature and at least one acousto-EM signature of the biologic.

Claims 8-10 meet the criteria set out in PCT Article 33(2)-(4), because the prior art does not teach or fairly suggest a method for targeting a biologic structure to affect at least one function which comprises irradiating the biologic structure with at least one electromagnetic (EM) property and/or field to result in acoustic energy having a frequency including at least one resonant acoustic frequency of the biologic structure, the acoustic energy being present in an amount sufficient to affect at least one function of the biologic structure.

Claims 11-15 meet the criteria set out in PCT Article 33(2)-(4), because the prior art does not teach or fairly suggest a method for targeting a biologic structure and affecting at least one function of the biologic structure by inducing acoustic resonance therein comprising applying at least two energies selected from the group consisting of at least one acoustic energy and at least one electromagnetic energy, wherein at least one and at least two energies result in biologic structure being in acoustic resonance and at least a second and at least two energies provides additional energy to biologic structure, and applying at least two energies such that a power intensity level is achieved to induce acoustic resonance within the targeted biologic structure and to affect at least one function thereof.

Claims 16-22 meet the criteria set out in PCT Article 33(2)-(4), because the prior art does not teach or fairly suggest a method for targeting a biologic structure to affect its function characterized by the step which comprises applying electromagnetic energy to the biologic structure to induce acoustic resonance therein and affect its functions.

Claim 23 meets the criteria set out in PCT Article 33(2)-(4), because the prior art does not teach or fairly suggest a method to induce acoustic stimulation of a biologic structure to detect and/or identify a biologic structure comprising: applying to the biologic structure at least one acoustic energy comprising at least one non-resonant frequency to stimulate the biologic structure; receiving at least one electromagnetic energy pattern from the structure; and determining at least one non-resonant electromagnetic signature of the stimulated biologic structure.

INTERNATIONAL PRELIMINARY EXAMINATION REPORT

International application No.
PCT/US99/20776

Supplemental Box

(To be used when the space in any of the preceding boxes is not sufficient)

Claim 24 meets the criteria set out in PCT Article 33(2)-(4), because the prior art does not teach or fairly suggest a system for inducing acoustic stimulation of a biologic structure to detect and/or identify a biologic structure comprising: means for applying to the biologic structure at least one acoustic energy comprising at least one non-resonant frequency to stimulate the biologic structure; means for receiving at least one electromagnetic energy pattern from the structure; and means for determining at least one non-resonant electromagnetic signature of the stimulated biologic structure.

Claims 25-28 meet the criteria set out in PCT Article 33(2)-(4), because the prior art does not teach or fairly suggest a method for detecting and/or identifying an inorganic or biologic structure characterized by the steps comprising inducing acoustic resonance in the structure and detecting at least one acousto-EM signature of the structure.

Claims 29 and 32-34 lack Novelty under PCT Article 33(2) as being anticipated by Tsuboi et al. (5,777,228). Tsuboi et al. discloses a system for identifying a structure by determining at least one resonant acoustic signature of the structure comprising: means for inducing acoustic resonance in the structure (42); means for detecting at least one acoustic signature of the structure (45), and means for comparing said at least one acoustic signature of the structure with at least one reference acoustic signature (50).

Claims 29 and 32-34 lack an Inventive Step under PCT Article 33(3) as being anticipated by Tsuboi et al.

Claims 30-31 meet the criteria set out in PCT Article 33(2)-(4), because the prior art does not teach or fairly suggest a system for identifying a structure by determining at least one resonant acoustic signature of the structure comprising: means for inducing acoustic resonance in the structure; means for detecting at least one acoustic signature of the structure; and means for comparing said at least one acoustic signature of the structure with at least one reference acoustic signature wherein said system further comprises a means for detecting at least one acoustic-EM signature of the structure.

Claim 35 meets the criteria set out in PCT Article 33(2)-(4), because the prior art does not teach or fairly suggest a system for inducing acoustic resonance in a biologic structure to augment at least one function of the biologic structure comprising: means for generating at least one acoustic signal; means for transmitting said at least one acoustic signal to the biologic structure; and means for controlling the power level of said at least one acoustic signal to augment at least one function of the biologic structure.

Claim 36 meets the criteria set out in PCT Article 33(2)-(4), because the prior art does not teach or fairly suggest a system for inducing acoustic resonance in a biologic structure to affect at least one function of the biologic structure comprising means for generating at least one electromagnetic, and means for transmitting said at least one electromagnetic signal to the biological structure.

Claim 37 meets the criteria set out in PCT Article 33(2)-(4), because the prior art does not teach or fairly suggest a system for determining induction of acoustic resonance in a structure comprising means for generating electromagnetic energy corresponding to at least one acousto-EM signature, means for transmitting said electromagnetic energy to the structure, means for receiving at least one signal from the structure when said electromagnetic energy has interacted with the structure, and means for determining induction of acoustic resonance in the structure.

Claim 38 meets the criteria set out in PCT Article 33(2)-(4), because the prior art does not teach or fairly suggest a method for determining induction of acoustic resonance in a structure characterized by the steps comprising irradiating the structure with electromagnetic energy corresponding to at least one acousto-EM signature, receiving at least one signal from the structure when electromagnetic energy has interacted with the structure, and determining induction of acoustic resonance in the structure.

Claim 39 meets the criteria set out in PCT Article 33(2)-(4), because the prior art does not teach or fairly suggest a method to affect at least one function of a living transducer biologic structure comprising applying at least one electromagnetic energy to the biologic structure, at least one electromagnetic energy comprising at least one frequency which includes at least one resonant frequency of the biologic structure to induce acoustic resonance within the biologic structure, the energy being present in an amount sufficient to affect at least one function of the biologic structure.

Claims 40-42 meet the criteria set out in PCT Article 33(2)-(4), because the prior art does not teach or fairly suggest a method for targeting an inorganic structure to affect structure, the method comprising applying at least one resonant acousto-EM energy.

Claim 43 meets the criteria set out in PCT Article 33(2)-(4), because the prior art does not teach or fairly suggest a method for detecting an inorganic structure comprising inducing acoustic resonance in the structure and detecting at least one resonant acousto-EM energy.

Claim 44 meets the criteria set out in PCT Article 33(2)-(4), because the prior art does not teach or fairly suggest a method for detecting an inorganic structure comprising: inducing acoustic resonance in the structure by applying an acousto-EM signature and detecting at least one acoustic signature.

INTERNATIONAL PRELIMINARY EXAMINATION REPORT

International application No.
PCT/US99/20776

Supplemental Box

(To be used when the space in any of the preceding boxes is not sufficient)

Claims 29 and 32-34 meet the criteria for Industrial Applicability set out in PCT Article 33(4), because the present claimed invention is useful in the industry.

----- NEW CITATIONS -----

NONE

PATENT COOPERATION TREATY

From the
INTERNATIONAL PRELIMINARY EXAMINING AUTHORITY

To:
MARK G. MORTENSON
THE LAW OFFICES OF MARK G. MORTENSON
P. O. BOX 310
NORTH EAST MD 21901-0310

PCT NOTIFICATION OF TRANSMITTAL OF INTERNATIONAL PRELIMINARY EXAMINATION REPORT

(PCT Rule 71.1)

		Date of Mailing (day/month/year)	29 DEC 2000
Applicant's or agent's file reference		IMPORTANT NOTIFICATION	
BLP:106 PCT			
International application No.	International filing date (day/month/year)	Priority date (day/month/year)	
PCT/US99/20776	10 September 1999 (10.09.1999)	11 September 1998 (11.09.1998)	
Applicant			
BERKSHIRE LABORATORIES, INC.			

1. The applicant is hereby notified that this International Preliminary Examining Authority transmits herewith the international preliminary examination report and its annexes, if any, established on the international application.
2. A copy of the report and its annexes, if any, is being transmitted to the International Bureau for communication to all the elected Offices.
3. Where required by any of the elected Offices, the International Bureau will prepare an English translation of the report (but not of any annexes) and will transmit such translation to those Offices.
4. REMINDER

The applicant must enter the national phase before each elected Office by performing certain acts (filing translations and paying national fees) within 30 months from the priority date (or later in some Offices)(Article 39(1))(see also the reminder sent by the International Bureau with Form PCT/IB/301).

Where a translation of the international application must be furnished to an elected Office, that translation must contain a translation of any annexes to the international preliminary examination report. It is the applicant's responsibility to prepare and furnish such translation directly to each elected Office concerned.

For further details on the applicable time limits and requirements of the elected Offices, see Volume II of the PCT Applicant's Guide.

Name and mailing address of the IPEA/US Commissioner of Patents and Trademarks Box PCT Washington, D.C. 20231 Facsimile No. (703) 305-3230	Authorized officer Hezron Williams Telephone No. (703) 308-0956 <i>Hezron Williams</i>
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Form PCT/IPEA/416 (July 1992)

14 09/785794

PATENT COOPERATION TREATY

PCT

REC'D 05 JAN 2001

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PCT

INTERNATIONAL PRELIMINARY EXAMINATION REPORT

(PCT Article 36 and Rule 70)

Applicant's or agent's file reference BLP:106 PCT	FOR FURTHER ACTION See Notification of Transmittal of International Preliminary Examination Report (Form PCT/IPEA/416)	
International application No. PCT/US99/20776	International filing date (day/month/year) 10 September 1999 (10.09.1999)	Priority date (day/month/year) 11 September 1998 (11.09.1998)
International Patent Classification (IPC) or national classification and IPC IPC(7): A61B 17/22, 6/00; A61N 7/00 and US Cl.: 600/427, 429, 439; 601/2, 3, 4; 73/579		
Applicant BERKSHIRE LABORATORIES, INC.		

1. This international preliminary examination report has been prepared by this International Preliminary Examining Authority and is transmitted to the applicant according to Article 36.
2. This REPORT consists of a total of 6 sheets, including this cover sheet.

☒ This report is also accompanied by ANNEXES, i.e., sheets of the description, claims and/or drawings which have been amended and are the basis for this report and/or sheets containing rectifications made before this Authority (see Rule 70.16 and Section 607 of the Administrative Instructions under the PCT).

These annexes consist of a total of 24 sheets.

3. This report contains indications relating to the following items:

- I ☒ Basis of the report
- II ☐ Priority
- III ☐ Non-establishment of report with regard to novelty, inventive step and industrial applicability
- IV ☐ Lack of unity of invention
- V ☒ Reasoned statement under Article 35(2) with regard to novelty, inventive step or industrial applicability; citations and explanations supporting such statement
- VI ☐ Certain documents cited
- VII ☐ Certain defects in the international application
- VIII ☐ Certain observations on the international application

RECEIVED
JAN 15 2001
TECHNOLOGY CENTER 2100

Date of submission of the demand 10 April 2000 (10.04.2000)	Date of completion of this report 11 December 2000 (11.12.2000)
Name and mailing address of the IPEA/US Commissioner of Patents and Trademarks Box PCT Washington, D.C. 20231 Facsimile No. (703) 305-3230	Authorized officer Hezron Williams Telephone No. (703) 308-0956 <i>Hezron Williams</i>

Form PCT/IPEA/409 (cover sheet)(July 1998)

INTERNATIONAL PRELIMINARY EXAMINATION REPORT

International application No.

PCT/US99/20776

I. Basis of the report

1. With regard to the elements of the international application:*



the international application as originally filed.



the description:

pages 2, 14-21, 23-25, 27-37, 39-40, 42-46, 49-53, 55-62, 64-66 as originally filedpages NONE, filed with the demandpages 1, 3-13, 22, 26, 38, 41, 47-48, 54 and 63, filed with the letter of 22 September 2000 (22.09.2000)

the claims:

pages NONE, as originally filedpages NONE, as amended (together with any statement) under Article 19pages NONE, filed with the demandpages 67-73, filed with the letter of 22 September 2000 (22.09.2000)

the drawings:

pages 1-28, as originally filedpages NONE, filed with the demandpages NONE, filed with the letter of _____.

the sequence listing part of the description:

pages NONE, as originally filedpages NONE, filed with the demandpages NONE, filed with the letter of _____.2. With regard to the **language**, all the elements marked above were available or furnished to this Authority in the language in which the international application was filed, unless otherwise indicated under this item.

These elements were available or furnished to this Authority in the following language _____ which is:



the language of a translation furnished for the purposes of international search (under Rule 23.1(b)).



the language of publication of the international application (under Rule 48.3(b)).



the language of the translation furnished for the purposes of international preliminary examination (under Rules 55.2 and/or 55.3).

3. With regard to any **nucleotide and/or amino acid sequence** disclosed in the international application, the international preliminary examination was carried out on the basis of the sequence listing:

contained in the international application in printed form.



filed together with the international application in computer readable form.



furnished subsequently to this Authority in written form.



furnished subsequently to this Authority in computer readable form.



The statement that the subsequently furnished written sequence listing does not go beyond the disclosure in the international application as filed has been furnished.



The statement that the information recorded in computer readable form is identical to the written sequence listing has been furnished.

4. ☒ The amendments have resulted in the cancellation of:the description, pages NONEthe claims, Nos. NONEthe drawings, sheets/fig NONE5. ☐ This report has been established as if (some of) the amendments had not been made, since they have been considered to go beyond the disclosure as filed, as indicated in the Supplemental Box (Rule 70.2(c)).**

* Replacement sheets which have been furnished to the receiving Office in response to an invitation under Article 14 are referred to in this report as "originally filed" and are not annexed to this report since they do not contain amendments (Rules 70.16 and 70.17).

** Any replacement sheet containing such amendments must be referred to under item 1 and annexed to this report.

INTERNATIONAL PRELIMINARY EXAMINATION REPORT

International application No.

PCT/US99/20776

V. Reasoned statement under Article 35(2) with regard to novelty, inventive step or industrial applicability; citations and explanations supporting such statement

1. STATEMENT

Novelty (N)	Claims <u>1-28, 30-31, 35-44</u>	YES
	Claims <u>29, 32-34</u>	NO
Inventive Step (IS)	Claims <u>1-28, 30-31, 35-44</u>	YES
	Claims <u>29, 32-34</u>	NO
Industrial Applicability (IA)	Claims <u>1-44</u>	YES
	Claims <u>NONE</u>	NO

2. CITATIONS AND EXPLANATIONS (Rule 70.7)

Please See Continuation Sheet

INTERNATIONAL PRELIMINARY EXAMINATION REPORT

International application No.
PCT/US99/20776

Supplemental Box

(To be used when the space in any of the preceding boxes is not sufficient)

V. 2. Citations and Explanations:

Claims 1-2 and 4-5 meet the criteria set out in PCT Article 33(2)-(4), because the prior art does not teach or fairly suggest a method for targeting a biologic structure to augment its function which comprises irradiating the biologic structure with acoustic energy to induce acoustic resonance therein.

Claims 3 and 6-7 meet the criteria set out in PCT Article 33(2)-(4), because the prior art does not teach or fairly suggest a method for targeting a biologic structure to affect at least one function which comprises irradiating the biologic structure with acoustic energy having a frequency near or at the resonant frequency of the biologic structure to induce acoustic resonance therein, and determining at least one acoustic signature and at least one acousto-EM signature of the biologic.

Claims 8-10 meet the criteria set out in PCT Article 33(2)-(4), because the prior art does not teach or fairly suggest a method for targeting a biologic structure to affect at least one function which comprises irradiating the biologic structure with at least one electromagnetic (EM) property and/or field to result in acoustic energy having a frequency including at least one resonant acoustic frequency of the biologic structure, the acoustic energy being present in an amount sufficient to affect at least one function of the biologic structure.

Claims 11-15 meet the criteria set out in PCT Article 33(2)-(4), because the prior art does not teach or fairly suggest a method for targeting a biologic structure and affecting at least one function of the biologic structure by inducing acoustic resonance therein comprising applying at least two energies selected from the group consisting of at least one acoustic energy and at least one electromagnetic energy, wherein at least one and at least two energies result in biologic structure being in acoustic resonance and at least a second and at least two energies provides additional energy to biologic structure, and applying at least two energies such that a power intensity level is achieved to induce acoustic resonance within the targeted biologic structure and to affect at least one function thereof.

Claims 16-22 meet the criteria set out in PCT Article 33(2)-(4), because the prior art does not teach or fairly suggest a method for targeting a biologic structure to affect its function characterized by the step which comprises applying electromagnetic energy to the biologic structure to induce acoustic resonance therein and affect its functions.

Claim 23 meets the criteria set out in PCT Article 33(2)-(4), because the prior art does not teach or fairly suggest a method to induce acoustic stimulation of a biologic structure to detect and/or identify a biologic structure comprising: applying to the biologic structure at least one acoustic energy comprising at least one non-resonant frequency to stimulate the biologic structure; receiving at least one electromagnetic energy pattern from the structure; and determining at least one non-resonant electromagnetic signature of the stimulated biologic structure.

INTERNATIONAL PRELIMINARY EXAMINATION REPORT

International application No.
PCT/US99/20776**Supplemental Box**

(To be used when the space in any of the preceding boxes is not sufficient)

Claim 24 meets the criteria set out in PCT Article 33(2)-(4), because the prior art does not teach or fairly suggest a system for inducing acoustic stimulation of a biologic structure to detect and/or identify a biologic structure comprising: means for applying to the biologic structure at least one acoustic energy comprising at least one non-resonant frequency to stimulate the biologic structure; means for receiving at least one electromagnetic energy pattern from the structure; and means for determining at least one non-resonant electromagnetic signature of the stimulated biologic structure.

Claims 25-28 meet the criteria set out in PCT Article 33(2)-(4), because the prior art does not teach or fairly suggest a method for detecting and/or identifying an inorganic or biologic structure characterized by the steps comprising inducing acoustic resonance in the structure and detecting at least one acousto-EM signature of the structure.

Claims 29 and 32-34 lack Novelty under PCT Article 33(2) as being anticipated by Tsuboi et al. (5,777,228). Tsuboi et al. discloses a system for identifying a structure by determining at least one resonant acoustic signature of the structure comprising: means for inducing acoustic resonance in the structure (42); means for detecting at least one acoustic signature of the structure (45), and means for comparing said at least one acoustic signature of the structure with at least one reference acoustic signature (50).

Claims 29 and 32-34 lack an Inventive Step under PCT Article 33(3) as being anticipated by Tsuboi et al.

Claims 30-31 meet the criteria set out in PCT Article 33(2)-(4), because the prior art does not teach or fairly suggest a system for identifying a structure by determining at least one resonant acoustic signature of the structure comprising: means for inducing acoustic resonance in the structure; means for detecting at least one acoustic signature of the structure; and means for comparing said at least one acoustic signature of the structure with at least one reference acoustic signature wherein said system further comprises a means for detecting at least one acoustic-EM signature of the structure.

Claim 35 meets the criteria set out in PCT Article 33(2)-(4), because the prior art does not teach or fairly suggest a system for inducing acoustic resonance in a biologic structure to augment at least one function of the biologic structure comprising: means for generating at least one acoustic signal; means for transmitting said at least one acoustic signal to the biologic structure; and means for controlling the power level of said at least one acoustic signal to augment at least one function of the biologic structure.

Claim 36 meets the criteria set out in PCT Article 33(2)-(4), because the prior art does not teach or fairly suggest a system for inducing acoustic resonance in a biologic structure to affect at least one function of the biologic structure comprising means for generating at least one electromagnetic, and means for transmitting said at least one electromagnetic signal to the biological structure.

Claim 37 meets the criteria set out in PCT Article 33(2)-(4), because the prior art does not teach or fairly suggest a system for determining induction of acoustic resonance in a structure comprising means for generating electromagnetic energy corresponding to at least one acousto-EM signature, means for transmitting said electromagnetic energy to the structure, means for receiving at least one signal from the structure when said electromagnetic energy has interacted with the structure, and means for determining induction of acoustic resonance in the structure.

Claim 38 meets the criteria set out in PCT Article 33(2)-(4), because the prior art does not teach or fairly suggest a method for determining induction of acoustic resonance in a structure characterized by the steps comprising irradiating the structure with electromagnetic energy corresponding to at least one acousto-EM signature, receiving at least one signal from the structure when electromagnetic energy has interacted with the structure, and determining induction of acoustic resonance in the structure.

Claim 39 meets the criteria set out in PCT Article 33(2)-(4), because the prior art does not teach or fairly suggest a method to affect at least one function of a living transducer biologic structure comprising applying at least one electromagnetic energy to the biologic structure, at least one electromagnetic energy comprising at least one frequency which includes at least one resonant frequency of the biologic structure to induce acoustic resonance within the biologic structure, the energy being present in an amount sufficient to affect at least one function of the biologic structure.

Claims 40-42 meet the criteria set out in PCT Article 33(2)-(4), because the prior art does not teach or fairly suggest a method for targeting an inorganic structure to affect structure, the method comprising applying at least one resonant acousto-EM energy.

Claim 43 meets the criteria set out in PCT Article 33(2)-(4), because the prior art does not teach or fairly suggest a method for detecting an inorganic structure comprising inducing acoustic resonance in the structure and detecting at least one resonant acousto-EM energy.

Claim 44 meets the criteria set out in PCT Article 33(2)-(4), because the prior art does not teach or fairly suggest a method for detecting an inorganic structure comprising: inducing acoustic resonance in the structure by applying an acousto-EM signature and detecting at least one acoustic signature.

INTERNATIONAL PRELIMINARY EXAMINATION REPORT

International application No.
PCT/US99/20776

Supplemental Box

(To be used when the space in any of the preceding boxes is not sufficient)

Claims 29 and 32-34 meet the criteria for Industrial Applicability set out in PCT Article 33(4), because the present claimed invention is useful in the industry.

----- NEW CITATIONS -----

NONE

**METHODS FOR USING RESONANT ACOUSTIC AND/OR RESONANT
ACOUSTO-EM ENERGY TO DETECT AND/OR EFFECT STRUCTURES**

TECHNICAL FIELD

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The present invention relates to detection of inorganic and biologic structures and/or disruption and/or augmentation of functions of structures using acoustic, resonant acoustic, and/or resonant acousto-EM energy and/or electromagnetic properties and/or fields.

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BACKGROUND OF THE INVENTION

The resonant acoustic frequency of a system is the natural free oscillation frequency of the system. A resonant acoustic system can be excited by a weak mechanical or acoustic driving force in a narrow band of frequencies, close or equal to the resonant frequency thereby inducing acoustic resonance in a targeted structure.

15

Acoustic resonance has been used to determine various properties of solid materials. For instance, Migliori et al in U.S. Patent Nos. 4,976,148 and 5,062,296 and 5,355,731 disclose a method for characterizing a unique resonant frequency spectroscopic signature for objects derived from ultrasonic excitation of objects, the use of resonant ultrasound spectroscopy for grading production quantities of spherical objects such as roller balls for bearings, and the use of resonant ultrasound spectroscopy with a rectangular parallelepiped sample of a high dissipation material to enable low amplitude resonance to be detected for use in calculating the elastic constants of the high dissipation sample. However, the Migliori patents are directed to solid materials and not to selectively targeting organic or biologic material especially when liquid systems are involved.

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In addition to interacting with inanimate structures, acoustic energy also interacts with living, biologic organisms and structures. Acoustic energy has been used extensively in medicine and biology for imaging structures, by directing an acoustic wave at a biologic structure and analyzing the reflection pattern of the acoustic wave. Also, acoustic energy has been used in physical therapy medicine for delivering heat to targeted areas of injury or pain. However, all of the above applications depend on using acoustic energy that is non-selective

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physical structure vibrates and the vibrational energy of motion may be transferred to the surrounding medium which includes air, liquid, or solid.

"Detect" as used herein is defined as determining the presence or absence of a structure, and if present identifying the structure.

5 **"Electromagnetic (EM) properties and/or fields"** as used herein includes direct and alternating currents, electric and magnetic fields, electromagnetic radiation, and fields which include but are not limited to waves, current, flux, resistance, potential, radiation or any physical phenomena including those obtainable or derivable from the Maxwell equations, incorporated by reference herein.

10 **"Electromagnetic (EM) energy pattern"** as used herein represents the electromagnetic energy produced by a structure as acoustic energy interacts with the structure and is manifested as electromagnetic properties and/or fields.

15 **"Biologic structure"** as used herein, and used interchangeably with organic, includes anything from the smallest organic or biochemical ion or molecule, to cells, organs, and entire organisms.

"Disruption" as used herein refers to deleterious effects on a structure.

"Acoustic signature" as used herein means a unique acoustic pattern that is produced by the structure when in acoustic resonance that may take the form of amplitude of signal.

20 **"Resonant acoustic frequency"** as used herein includes frequencies near or at the natural resonant frequency of the structure including harmonic and subharmonic frequencies of the natural resonant frequency to induce acoustic resonance therein.

25 **"Acousto-EM signature"** as used herein is defined as an EM energy pattern of an object in acoustic resonance and/or an EM energy equivalent in frequency to the resonant acoustic frequency.

"Acousto-EM spectroscopy" as used herein is defined as detecting a unique EM signature for a structure that is in acoustic resonance, or detecting a unique acoustic signature from a structure that is in resonance due to the introduction of electromagnetic energy, both of which can be used to detect and/or identify the structure in resonance.

30 **"Living transducer"** as used herein is defined as a biologic structure, such as a piezoelectric or semiconductor that converts electromagnetic energy or fields into mechanical

energy and/or mechanical energy into electromagnetic energy or fields.

"**Cavitation**" as described herein is defined as the formation of vapor-filled cavities in liquids, e.g., bubble formation in water when brought to a boil.

5 "**Mechanical**" as described herein include mechanisms such as compression and rarefaction which are thought to take place in the intensity/duration threshold region between the thermal and cavitation regions.

 "**Non-resonant electromagnetic signature**" as used herein is defined as an EM energy pattern produced by an object stimulated by a non-resonant acoustic field.

10 "**Resonant acousto-EM energy**" as described herein means electromagnetic properties and/or fields that induce acoustic resonance in a structure.

 The present invention addresses the shortcomings of the prior art by inducing acoustic resonance in a targeted structure with select frequencies that affect the specific targeted structure but have virtually no effect on nearby, non-resonating structures. Furthermore, acoustic energy power intensities can be reduced by introducing a source of
15 electromagnetic (EM) energy that augments, or replaces, the acoustic energy thereby reducing the destructive nature of high power acoustic energy. The interaction between EM energy and acoustic resonance allows for precise detection of a structure in acoustic resonance by producing a signature with high signal to noise ratio, while producing little effect in other structures.

20 The present invention provides methods to selectively detect, identify and/or affect an inorganic or biologic structure by using resonant acoustic and/or acousto-EM energy which can transfer useful energy to targeted structures while leaving nearby structures, which are not in resonance, virtually unchanged.

 Therefore, it is an object of the present invention to provide a method of identifying
25 or detecting an inorganic or biologic structure using its resonant acoustic and/or acousto-EM signature and/or EM energy patterns.

 It is an object of the present invention to provide a method for using resonant acoustic and/or acousto-EM signatures and/or energy patterns to augment and/or disrupt the growth and/or function of biologic structures.

30 It is another object of the invention to provide a method for determining resonant frequencies of a biologic structure.

 It is also an object of the invention to provide a method using resonant acoustic

and/or resonant acousto-EM energies to detect the presence of and/or identify biologic structures.

5 In accordance with the aforesaid objects the present invention provides for the detection of inorganic or biologic structures and/or disruption and/or augmentation of growth and/or functions of said structures using resonant acoustic and/or resonant acousto-EM signatures and/or EM energy patterns.

Applying principles of acoustic resonance, the resonant acoustic frequency of a biologic system is the natural free oscillation frequency of the system, and thus a can be excited by a weak mechanical or acoustic driving force in a narrow band of frequencies.

10 Also, depending on the size, shape, and composition of the biologic structure, there can be more than one naturally occurring resonant acoustic frequency, as well as numerous subharmonic and superharmonic resonant acoustic frequencies.

When a structure, including both inorganic and biologic structures, goes into acoustic resonance, energy builds up in it rapidly. The energy is either kept in the system or released

15 to the surrounding environment. Energy kept in the structure can enhance the structure's functions or cause disruption of the structure. The energy in a resonant system is either intrinsically dissipated as electromagnetic energy and/or is transmitted as acoustic energy to the nearby medium. The intrinsically dissipated energy is of particular interest, because it is dissipated through molecular and atomic vibrations, producing EM energy patterns. This EM

20 energy is referred to as acousto-EM energy because it is produced when a structure is excited by acoustic energy and some acoustic energy interacts with the structure and is converted into electromagnetic energy thereby being intrinsically dissipated. The properties, fields and/or frequencies of EM energy produced depend on the unique molecular and atomic components of the structure in question. Moreover, the induction of acoustic resonance in

25 a structure leads to the production of a unique acousto-EM signature for that structure, which can be used to detect and/or identify the structure as disclosed in the present invention. Conversely, if a structure is targeted with an applied EM energy equivalent to its acousto-EM signature, the energy dissipation pathway is reversed, and a state of acoustic resonance can be induced. Reversing the energy dissipation pathway with an applied acousto-EM signature can be

30 used to produce the same augmentation, detection, and disruption effects that the original resonant acoustic energy field produces. An applied acousto-EM signature can be used either by itself, or in

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combination with resonant acoustic energy. Using the applied resonant acousto-EM signature and resonant acoustic energy together, allows for the use of lower power levels of both types of energy, lessening the potential adverse affects of electromagnetic energy and/or acoustic energy on nearby or adjacent nontargeted structures.

- 5 Electromagnetic energy may also interact with and complement an acoustic energy wave in a system in at least four ways: via the piezoelectric effect, intrinsic dissipation of electromagnetic energy and via the acoustoelectric or magnetoacoustic effect.

- 10 In the piezoelectric effect, acoustic vibratory energy is converted interchangeably with EM energy by a transducer. Biologic piezoelectric structures can modulate the same conversion of energy, thereby acting as living transducers. Thus, when an EM field is applied to a biologic piezoelectric structure, an acoustic wave is produced. Likewise, when an acoustic wave is applied to a biologic piezoelectric structure, EM energy is produced. The piezoelectric effect in biologic structures has many useful applications (see below.) This effect becomes even more useful when principles of acoustic resonance are applied. In the present invention specific biologic structures can be targeted with an acoustic wave or EM energy at power levels that dramatically affect the target structure, but have virtually no effect on adjacent, nonresonant structures. Although not previously postulated by others, biologic structures functioning as living, resonant piezoelectric transducers which modulate the conversion of mechanical and EM energy is undoubtedly one of the major underlying mechanisms responsible for the interaction of EM fields with biologic structures.
- 15 In the acoustoelectric effect, the passage of an acoustic wave through a semiconductor induces an electric current. The passage of an acoustic wave through the material is postulated to cause a periodic spatial variation of the potential energy of the charge carriers. This results in an electric field across the ends of the semiconductor as long as the acoustic wave is traversing the semiconductor. Free electron carriers are bunched in the potential-energy troughs, and as the acoustic wave having a specific frequency propagates, it drags the bunches along with it, resulting in an electric field such as a DC field pulsing at the specific acoustic frequency or an AC field having a frequency equal to the specific acoustic frequency. The effect is enhanced where there are both positively and negatively charged carriers, and where there are many different groups of carriers - conditions which are frequently found in biologic systems. The attributes of the current
- 20
- 25
- 30

produced depend on the unique molecular and atomic components of the structure in question. This aspect alone provides a means to perform acoustoelectric spectroscopy on biologics many of which are semiconductors, and depending on the selected frequency, the acoustoelectric effect in biological structures has many other potentially useful applications.

5 Thus understood, a targeted structure can be irradiated or exposed to acoustic energy having non-resonant frequency and an electromagnetic energy pattern of the acoustoelectric effect in the structure can be detected. This detected non-resonant electromagnetic signature can be used as a signature to affect, detect and identify the targeted structure.

10 However, the acoustoelectric effect becomes even more useful when principles of acoustic resonance are applied. Augmentation, detection, and/or disruption of biologics can be targeted to specific structures at power levels that dramatically affect the target structure, but have virtually no effect on nearby, nonresonant structures. The current produced by the acoustoelectric effect in a resonant structure will be much stronger than any current produced by neighboring non-resonant structures, and may be of an alternating nature. The large signal
 15 to noise ratio obtained from a resonant structure improves accuracy of acoustic and EM energy pattern identification and detection. Similar to reversal of the piezoelectric effect and acoustic resonance intrinsic energy dissipation pathway (see above), application of the resonant acoustoelectric EM energy pattern to a targeted structure will amplify the acoustic wave (acoustoelectric gain which peaks at the frequency for which the acoustic wavelength
 20 is the Debye length, where bunching is optimum). Thus, combined use of the resonant acoustic, acoustoelectric and/or EM fields permit greater tissue penetration of high frequency acoustic energy that would otherwise be highly attenuated and have poor tissue penetration. Using the resonant acoustic frequency, acoustoelectric and/or EM fields together also allows for the use of lower power levels of these types of energy, lessening the potential effects on
 25 other nontargeted and nonresonant structures.

The magnetoacoustic effect is the magnetic-field-dependent attenuation of an acoustic field in a monotonic, oscillatory, or resonant manner, depending on the electronic properties of the substance in question. This variability in result, depending on structural composition, provides a further enhancement of acousto-EM spectroscopy in relation to
 30 biologics and other structures, via addition of a magnetic field. Also, the addition of a magnetic field provides the means to amplify or attenuate an acoustic field, thus improving or modulating

the penetration of the acoustic field in biologic tissues.

Similarly, resonant acoustics combined with acoustic cyclotron resonance (ie. resonant acoustic cyclotron resonance) and Doppler-shifted resonant acoustic cyclotron resonance presents a powerful, and precise means of selectively causing augmentation,
5 detection and/or disruption of structures.

The present invention provides a method that applies the principles of acoustic resonance to biologic structures for the purpose of disruption and/or augmentation of functions of the specifically targeted biologic structure. The resonant acoustic frequency of a biologic structure may be determined by performing resonant acoustic spectroscopy using
10 methods and systems well know in the art. Particularly, a resonant acoustic frequency of a biologic structure may be determined by the steps of:

- a) applying acoustic energy to the biologic structure and scanning through a range of acoustic energy frequencies; and
- b) detecting at least one specific frequency which causes a maximum signal output
15 from the biologic structure indicating the biologic structure being induced into acoustic resonance by the at least one specific frequency.

The specific frequencies causing the maximum signals are the resonant acoustic frequencies of the biologic structure which are defined and used herein as the acoustic signature of the biologic structure. Once determined, at least one resonant acoustic
20 frequency may be applied to the biologic structure to affect functioning therein and/or to determine its acousto-EM signature.

The acoustic energy, including the resonant acoustic frequencies (i.e., the acousto-EM signature) may be applied at a power level sufficient to affect functioning of the biologic structure. Depending on the power intensity of the acoustic energy, and the type of targeted
25 structure that is induced into acoustic resonance, the structure may have its functions affected, such as disruption and/or augmentation.

At lower power levels functions of the biologic structure can be augmented while at higher power levels disruption of the structure may occur. Augmentation as used herein encompasses beneficial effects on the biologic structure. Such augmenting of functions or
30 enhancing effects include but are not limited to enhancement of growth, reproduction, regeneration, embryogenesis, metabolism, fermentation, and the like. The results of such

enhancement include but are not limited to increase in bone mass or density, increase in number and maturation of eggs, increase in number and/or function of leukocytes, increase in fermentation products in beer, wine and cheese manufacturing, increase in plant germination and growth and the like.

5 There are some situations where the ability to selectively disrupt a structure with acoustic resonance is very useful as disclosed in the present invention. As stated above, disruption as used herein refers to deleterious effects on the biologic structure. Such deleterious effects include but are not limited to structural failure of the biologic structure resulting in lysis, shattering, rupture or inactivation of the biologic or of one or more
10 components of the biologic structure. Disruption as used herein also includes within its ambit inhibition of vital processes required for growth, reproduction, metabolism, infectivity and the like. Components which may be targeted for disruption include, but are not limited to DNA, RNA, proteins, carbohydrates, lipids, lipopolysaccharides, glycolipids, glycoproteins, proteoglycans, chloroplasts, mitochondria, endoplasmic reticulum, cells, organs and the like.
15 In the case of virulent organisms, the virulence factors may be specifically targeted for disruption to prevent or inhibit the growth, infectivity or virulence of the organism. Such virulence factors include but are not limited to endotoxins, exotoxins, pili, flagella, proteases, ligands for host cell receptors, capsules, cell walls, spores, chitin, and the like.

 Organics, biologics or one or more targeted portions thereof which are amenable to
20 disruption using the methods of the present invention include but are not limited to viruses, bacteria, protozoans, parasites, fungi, worms, mollusks, arthropods, tissue masses, and the like. The organics or biologics to be disrupted may be isolated, present in a multicellular organism or portion thereof, or other complex environment.

 It is postulated that disruption of the targeted biologic structure without affecting
25 nearby tissue or structures occurs due to acoustic resonance being induced only in the targeted structure which until now has not been considered a mechanism to affect a biologic structure. This is very different from that disclosed in the prior art which contemplates only three mechanisms for affecting a biologic structure which include cavitation, thermal and mechanical.

30 At specific power levels, such as in lower levels, that do not cause the actual disruption of a structure, resonant acoustic energy can intrinsically dissipate within the

structure. This intrinsically dissipated acoustic energy can be converted by the structure into an electromagnetic energy having specific properties and/or fields that may be manifested as direct and alternating currents, electric and magnetic fields, electromagnetic radiation and the like. The pattern of the electromagnetic energy represents a produced acousto-EM signature
5 of the structure.

The present invention provides a method to determine an acousto-EM signature of a structure which comprises irradiating the structure with acoustic energy having a frequency at or near a previously determined resonant acoustic frequency of the structure to induce resonance therein and detecting the electromagnetic energy pattern caused by the intrinsic
10 dissipation of energy.

Once an acousto-EM signature is determined for a specific structure, this structure can be induced into acoustic resonance by applying an EM energy pattern or equivalent to the acousto-EM signature of the structure. Typical electromagnetic energies applied include direct and alternating current, electric and magnetic fields, and electromagnetic radiation and
15 the like.

As such, the present invention applies the principles of acoustic resonance by applying resonant acoustic frequencies and electromagnetic energy equivalent to the predetermined acousto-EM signature of a targeted structure individually, or in combination, to affect the targeted structure, the method comprising the steps of:

- 20 a) applying at least one resonant acoustic frequency of the targeted structure; and/or
 b) applying electromagnetic energy equivalent to part or all of the acousto-EM signature of the targeted structure; and
 c) applying (a) and/or (b) each at a power intensity level to induce acoustic resonance within the targeted structure and affect functioning of the structure.

25 Either the resonant acoustic frequency of the targeted structure or the acousto-EM signature must be predetermined, as discussed above, to provide the applicable energy for inducing acoustic resonance in the structure. The electromagnetic energy can be introduced into the targeted structure in the form of a direct or alternating current having a specific frequency that is equivalent to the electromagnetic energy pattern (i.e., the acousto-EM signature)
30 detected when the structure is induced into acoustic resonance. Furthermore each type of energy can be applied at a power level less than used individually and this allows for inducing acoustic resonance

in the structure with the possibility of reducing damage to the structure.

The present invention provides a method for detecting and/or identifying inorganic or biologic structures using resonant acoustic and/or acousto-EM energy. The method includes determining the acoustic signature of a structure by irradiating the structure with a
5 range of frequencies to determine the specific frequency and/or frequencies that induce acoustic resonance therein to provide an acoustic signature of the structure. The acoustic signature can be compared with reference signatures to detect and/or identify the structure.

Furthermore, the identification and/or detection of a structure can also be achieved by detecting an acousto-EM signature of a targeted structure, the method comprising the
10 steps of:

- a) inducing acoustic resonance in the targeted structure; and
- b) detecting an electromagnetic energy pattern from the targeted structure in
acoustic resonance which represents an acousto-EM signature of the structure.

The acousto-EM signature can be compared to reference signatures to detect and/or identify
15 the structure.

The targeted structure can be induced into acoustic resonance by introducing acoustic energy including at least one resonant acoustic frequency, electromagnetic energy equivalent to the resonant acoustic frequency, and/or an electromagnetic energy pattern equivalent to the acousto-EM signature.

20 The electromagnetic energy pattern manifested as electromagnetic properties and fields may be determined by detection means well known to those skilled in the art such as those disclosed in *Introduction to Electromagnetic Fields and Waves*, by Erik V. Bohn Addison-Wesley Publishing Co., 1968, the contents of which are incorporated by reference herein.

25 In another embodiment of the present invention, a structure may be induced into acoustic resonance by applying to the structure part or all of the acousto-EM signature of the structure to induce the structure into acoustic resonance. If the structure is induced into acoustic resonance, this fact may be used to detect and/or identify the structure. This represents another method of the present invention that may used for identification or
30 detection of a specific structure, because each structure will not only have its own unique acoustic signature but also will have a unique acousto-EM signature to which it responds by

resonating acoustically. Also, depending on the power intensity of the electromagnetic properties and/or fields and the type of targeted structure that is induced into acoustic resonance, the structure may have its functions affected, such as disruption and/or augmentation.

5 In all the above embodiments the introduction of acoustic and/or electromagnetic energy including a resonant acoustic frequency can be applied in either continuous and/or periodic form depending on the desired effect.

The acoustic and/or EM energy or fields may be applied individually or in combination. Likewise the acoustic and/or EM energy or fields may be detected individually or in
10 combination.

Many biochemical compounds and biologic structures are naturally occurring crystals and especially susceptible in that regard to the effects of resonant acoustic energy. Many biologic substances are piezoelectric materials. For instance, bone is a piezoelectric material and the piezoelectric properties of bone play a vital role in its biological functions. As such,
15 it is further envisioned by the inventors that biologic structures having a piezoelectric nature may be affected by applying a sufficient amount of acoustic energy and/or electromagnetic energy to induce the structure into resonance thereby affecting the functions of the biologic structure either positively or negatively. Thus understood, biologic structures that act as living transducers may be induced into acoustic resonance by introducing electromagnetic
20 energy equivalent to a resonant acoustic frequency of the biologic structure which is converted to mechanical energy by the living transducer thereby inducing acoustic resonance in the structure.

Another aspect of the invention is a system for detecting a biologic or inorganic structure by determining the resonant acoustic and/or acousto-EM signature of the structure
25 comprising:

- a) means for inducing acoustic resonance in the biologic or inorganic structure;
- b) means for detecting the acoustic signature of the biologic or inorganic structure; and
- c) means for comparing the acoustic signature of the biologic or inorganic structure
30 with a reference acoustic signature of the structure.

Also, the above system may also or instead comprise means for detecting a resonant acousto-EM energy signature of the structure in acoustic resonance which produces an

electromagnetic energy pattern such as described above. The acousto-EM signature can be compared with a previously determined reference acousto-EM signature by providing means for comparing in a detection or identification system. The electromagnetic energy pattern is manifested as electromagnetic properties and/or fields that include but are not limited to energy in the form of direct and alternating current, electric and magnetic fields, and electromagnetic radiation. The targeted structure can be induced into acoustic resonance by introducing acoustic energy including at least one resonant acoustic frequency, electromagnetic energy equivalent to the resonant acoustic frequency, and/or an electromagnetic energy pattern equivalent to the acousto-EM signature.

In another embodiment of the present invention a system for augmenting and/or disrupting a targeted biologic structure comprises means for applying acoustic energy including a previously determined resonant acoustic frequency to induce acoustic resonance in the biologic structure, the acoustic energy being applied at a sufficient power input to affect functions of the biologic structure. Alternatively, the targeted structure may be induced into acoustic resonance by providing electromagnetic energy equivalent to the resonant acoustic frequency or the acousto-EM signature that was previously determined, such electromagnetic energy including direct and alternating current, electric and magnetic fields, and electromagnetic energy.

In yet another embodiment a system is provided to introduce acoustic energies having acoustic frequencies at or near the resonant acoustic frequencies of the targeted structure and also electromagnetic energy to augment the resonant acoustic frequencies comprising:

means for introducing a frequency at or near the resonant acoustic frequency of the targeted structure ; and

means for introducing electromagnetic energy equivalent to the electromagnetic energy pattern previously determined as an acousto-EM signature of the structure, such means including direct and alternating current, electric and magnetic fields, and/or electromagnetic radiation and the like.

The acoustic energy and EM energy equivalent to the acousto-EM signature may be applied and/or detected by a single means that can apply both types of energy.

Additional objects, advantages and novel features of the invention will be set forth in part in the description which follows, and in part will become apparent to those skilled in

identical proteins.

In some viruses, a lipoprotein membrane, or envelope, surrounds the capsid. The envelope is derived from host cell membranes and is modified by the virus during its departure from the host cell. The envelope may carry specific virus proteins such as hemagglutinin or neuraminidase that are important for future functions and survival of the virus. The envelope of some viruses is studded with projections, or peplomers, which look like a fringe around the edge. The fringe may also be important for function and survival of the virus.

Classically, the piezoelectric phenomenon is said to exist when the application of a mechanical stress to certain dielectric (electrically nonconducting) crystals produces electric polarization (electric dipole moment per cubic meter) which is proportional to the mechanical stress. Conversely, application of an EM field to a crystal produces mechanical stress and distortion, and hence acoustic energy.

A necessary condition for the piezoelectric phenomenon in a crystal is the absence of a center of symmetry. Twenty of the 32 classically defined crystal classes lack a center of symmetry and are piezoelectric. Viruses are crystalline structures and as such are susceptible to vibrational effects by the use of resonant acoustic and/or acousto-EM energy. Icosahedral viruses have 5-fold symmetry and thus do not have a classical center of symmetry in their crystalline structure, the necessary condition for a piezoelectric substance. Helical viruses likewise do not have a classical center of symmetry, as the spiraling capsids are offset from the 90 degree horizontal of the center axis. In addition to the crystalline structure of viruses being susceptible to the vibrational resonant effects of acoustic energy, viruses, as used in the present invention, may also function as piezoelectric, acoustic resonance structures.

The classical 32 groups of naturally occurring crystals defined in non-organic chemistry, do not include a group with 5-fold or offset helical symmetry. It is postulated by the inventors that viruses may represent a 33rd and 34th group of naturally occurring crystals.

The present invention has the potential to significantly reduce the number and severity of viral infections suffered by the world population. The invention has the potential to augment production of vaccines, or viral gene transfer. Also, the present has veterinary

In another embodiment, receiving acoustic transducer mode also detects qualitative and quantitative resonant acoustic frequencies of the virus in the multicellular organism to determine efficacy and duration of treatment.

The present invention also provides a means to determine qualitative and quantitative resonant acoustic and/or acousto-EM frequencies *in vitro* as shown in Figure 19 A&B. A test device, as described above and shown in Figure 12, with any and all embodiments, is fitted with transmitters and receivers to transmit, detect, measure, and analyze EM energy. When the resonant acoustic frequencies are applied to the virus test disk, a unique electromagnetic energy pattern is generated, according to the structure and composition of the virus and test disk under study, referred to herein as the resonant acousto-EM signature. Mechanisms producing the resonant acousto-EM signature include, but are not limited to piezoelectricity, acoustoelectricity, magnetoacoustics, and/or intrinsic energy dissipation. The resonant acousto-EM signature represents one or more of several electromagnetic properties and/or fields including, but not limited to, direct current, alternating current, magnetic field, electric field, EM radiation, and/or acoustic cyclotron resonance (standard or Doppler shifted).

All of the above mentioned forms of EM energy are detected, measured, and analyzed with devices and methods known to those skilled in the art. (It should be noted that useful information may also be derived from application of nonresonant frequencies, ie. current characterization of semiconductor biologics via the acoustoelectric effect.) This data in combination with resonant signatures yields even greater sensitivity and specificity to the method. For example, Herpes simplex virus (HSV) I and II will have nearly identical resonant acoustic signatures because they are virtually identical in size and shape. They differ in molecular protein configuration, however, and can be distinguished by their acousto-EM signatures. This includes, but is not limited to, characterization at nonresonant and resonant frequencies of acoustoelectric currents, acousto-EM signatures produced via intrinsic energy dissipation, of acoustic modulation or attenuation in the presence of a magnetic field via the magnetoacoustic effect, and of electric or magnetic fields induced or affected by any of the above processes.

In another embodiment, the test device is also fitted with any and all combinations of resonant acoustic and acousto-EM generating equipment. A sample of unknown composition

a multicellular organism using a resonant acoustic and/or acousto-EM field probe. For example, as shown in Figure 23, a hand-held probe is fitted with an EM radiation generating device, as currently known to those skilled in the art. A predetermined EM radiation field (frequencies, harmonics, amplitude, mode, shape, etc.) replicating the acousto- EM signature representing the intrinsic dissipation pattern of a particular virus, is delivered to a predetermined portion of the organism, from the hand-held probe. For example, in a person afflicted with an upper respiratory tract infection (a "cold"), the treatment is delivered through the skin over the nose, throat, and sinuses, reversing the intrinsic energy dissipation pathway of the rhinovirus and inducing resonant acoustic oscillations which disrupt the rhinovirus.

Example 2.

Disruption, Augmentation, Detection and/or Identification of Micro-organisms

Any micro-organism, such as bacteria, as well as structure and molecules contained or associated herewith, may be augmented, disrupted, detected and/or identified *in vitro* or *in vivo* using the methods of the present invention. Bacteria include, but are not limited to, those associated with animals, man, avians, reptiles, amphibians, insects, aquatic life, plants, fruit, soil, water, oil, fermentation processes for food production, and the like. In one embodiment the bacteria include but are not limited to *Streptococcus* *sps.*, *Staphylococcus* *sps.*, *Hemophilus* *sps.*, *Neisseria* *sps.*, *Treponema* *sps.*, *Salmonella* *sps.*, *Shigella* *sps.*, *Escherichia coli* strains, *Corynebacteria* *sps.*, *Bordetella* *sps.*, *Chlostridium* *sps.*, *Rickettsia* *sps.*, *Chlamydia* *sps.*, *Brucella* *sps.*, *Mycobacterium* *sps.*, *Borrelia* *sps.*, *Mycoplasma* *sps.*, *Lactobacillus* *sps.*, strains thereof and the like. Human illnesses caused by bacteria include pneumonia, skin and wound infections, heart valve infections, gastroenteritis, syphilis, gonorrhea, the plague, urinary tract infections, lyme disease, tuberculosis, cholera, typhoid fever, anthrax, tetanus, and gangrene.

Fungal infections include athlete's foot, ringworm, vaginal yeast infections, oral thrush, histoplasmosis, and cryptococcus.

Diseases in animals caused by bacteria, fungi, protozoa and worms are similar to those in humans. Similarly, a wide range of micro-organisms infect plants, and even other micro-organisms are deemed to be beneficial (e.g., bakers yeast.).

Example 4

Augmentation of Bone Growth

Bone demineralization in humans is a significant health care problem. Thousands of
5 elderly people sustain fractures of the hip, leg, or arm due to this bone demineralization
(osteoporosis). These injuries cost the American health care system billions of dollars a year,
for treatment, surgery, and rehabilitation after the injury. In addition, the overall health status
of the victims is impaired, and they suffer loss of time and quality of life due to these
fractures. Other conditions which contribute to bone matrix loss include weightlessness (e.g.,
10 in outer space) and prolonged confinement to bed. People in certain occupations may benefit
from an increase in the normal bone density. Examples include professional athletes, military
personnel, and jobs requiring exposure to increased atmospheric pressures (e.g., undersea
diving).

Living bone is organized in a calcium based crystalline structure of hydroxyapatite,
15 doped with copper, and embedded in collagen fibers. The application of force to the collagen
fibers in the bony matrix, through mechanical pressure or gravitational fields, stimulates the
piezoelectric effect and flow of ions via fluid channels in bone. This small electrical charge,
in turn, acts as a signal to the body's osteoblasts to deposit more hydroxyapatite. As the
hydroxyapatite density increases, the bone becomes stronger. Thus, bones maintain their
20 normal structure and density in response to pressures and forces encountered in normal daily
activities, via a piezoelectric effect.

With aging, normal copper doping is lost, and the piezoelectric effect diminished.
The result is that hydroxyapatite density is not maintained, and the elderly suffer from
osteoporosis and bone fractures. The same thing occurs in the absence of normal activity
25 (weightlessness and confinement to bed), with subsequent absence of the normal piezoelectric
effect and ionic current flows.

Bone is a crystalline piezoelectric structure and as such is subject to the vibratory
effects of acoustic energy. The operative process behind normal physiologic bone density
maintenance is the generation of hydroxyapatite molecular movement within collagen fibers,
30 compressed by macro-pressures. These occur from daily activities, and stimulate the
piezoelectric and subsequent bone building osteoblastic effects.

This molecular movement and the collagen fiber compression can also be generated from micro-pressures within the semiconductor matrix of bone. Thus understood, micro-pressures can be produced by acoustic energy waves.

In addition to the piezoelectric effect, since bone is a piezoelectric and semiconductor structure, it will exhibit the acoustoelectric, intrinsic dissipation and magnetoacoustic effects. Conditions with diminished bone semiconductor function (osteoporosis) and/or decreased macro-pressures (weightlessness and bed confinement) can be effectively treated through application of acoustic micro-pressures which generate a biological piezoelectric effect, and/or also via acoustic resonance, intrinsic dissipation, acoustoelectric and magnetoacoustic effects.

Prior literature describes the use of non-resonant ultrasound to speed the rate of healing of bone fractures, however, the mechanism causes gross disruption of the bone tissues, which in turn damages the microscopic capillary bed in bone, with leakage of serum and cells into the bony matrix, and with subsequent bone mineralization. The literature also describes attempts to use ultrasound to detect resonant frequencies of the structure of entire bones (femur and ulna) to diagnose a bone as normal or defective. However, the use of resonant acoustics and/or acousto-EM frequencies to activate the piezoelectric effect is not described. No consideration is given in the prior art to using bone as a living transducer for the piezoelectric, intrinsic dissipation, acoustoelectric, and magnetoacoustic effects, either alone or in combination with a resonant acoustic field.

The present invention takes advantage of the crystalline, piezoelectric structure of bone for the purpose of augmenting bone growth and calcification. The invention has the potential to significantly reduce the number and severity of bone fractures suffered by victims of osteoporosis. The invention has the potential to speed the healing process of fractures. Other conditions which contribute to bone matrix loss, such as weightlessness (i.e., in outer space), or prolonged confinement to bed, would also benefit from the invention. The invention has the potential to aid people in occupations which would benefit from an increase in their bone density (athletes, military personnel, and jobs requiring exposure to increased atmospheric pressures such as undersea diving.) The invention also has potential veterinary applications. Unlike prior treatment using ultrasound, the present invention uses resonant acoustic and/or acousto-EM frequencies of bone to stimulate at least the piezoelectric effect

endangered species could be aided through the use and detection of resonant acoustic and/or acousto-EM frequencies specific for those organisms. The use of resonant acoustic and/or acousto-EM frequencies could potentially aid in the identification and differentiation of species and subspecies throughout the animal, plant, and microbiological kingdoms.

5 Examples of multicellular organisms whose growth and augmentation are desired for harvesting include plants and protein sources such as fish, clams, shrimp, chickens, and other livestock. Medicines, drugs, and chemicals harvested from a wide variety of plant and animal sources include hormones, perfumes, dyes, and vitamins. Other materials harvested from plant and animal sources are such an intrinsic part of human activities that they are simply too
10 numerous to list (i.e., pearls, clothing fibers, building materials, leather, etc.) At lower power inputs of the resonant acoustic and/or acousto-EM frequencies, these organisms and their structures can be selectively augmented.

 The present invention takes advantage of the discrete shape and size of numerous organisms to make use of resonant acoustic and/or acousto-EM frequencies specific to those
15 organisms, for purposes of augmentation, identification, detection and/or disruption. Using the piezoelectric, intrinsic energy dissipation, acoustoelectric, and/or magnetoacousto effects, the invention has the potential to produce the above results by applying an electromagnetic energy pattern of the specific acousto-EM signature, either alone or in combination with a resonant acoustic field. The present invention has the potential to provide chemical-free
20 control of numerous pests. The present invention also has the potential to provide for the detection and identification of numerous species of organisms. Lastly, the present invention has the potential to augment growth and metabolism in and of structures in various species deemed beneficial.

 The present invention provides a means to augment, detect, and/or disrupt
25 structures of multicellular organisms using resonant acoustic and/or acousto-EM energy. For example, as shown in Figure 32, a transducer apparatus with the resonant frequency for the cement plate of barnacles (by which they attach themselves to the hulls of ships) is fitted into an underwater "scrubber" which is operated remotely from the deck of the ship via cables, or from inside the vessel via RF control. As the scrubber moves along the outside of the hull,
30 the acoustic wave disrupts the cement plate of the barnacles, causing them to lose their grip on the hull and fall off into the ocean.

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DAY 3 - 11 of the peas exposed to the acousto-EM field sprouted while only 5 of the control peas sprouted. The acousto-EM exposed peas were almost twice the length of the control peas.

DAY 6 - 45 of the peas exposed to the acousto-EM field had sprouted while only 5 35 of the control group had sprouted.

DAY 10 - 61 of the peas exposed to the acousto-EM field had sprouted while only 45 of the control group had sprouted. The average length of the leaf sprout on the exposed acousto-EM field group was 3.3 cm while the average length of the control group was only 2.7 cm.

10 RESULTS: Applying an acousto-EM signature augmented the germination and growth rate of the peas.

Example 11

Detection and Identification of Inorganic Structures

The methods and systems of the present invention have a wide range of useful applications, such as on-site identification both qualitatively and quantitatively of various 15 types of inorganic matter or structures, recognition of impurities in metal alloys, recognition of armaments and weapons, such as plastic explosives, etc.

Detection and identification can be achieved by applying acoustic energy at a frequency closely matching the resonant frequency of an object or structure thereby inducing 20 acoustic resonance therein for detection of a unique acoustic and/or acousto-EM signature.

Using methods known to those skilled in the art, any device capable of generating and transmitting acoustic energy through any medium can be used to generate the resonant acoustic and/or acousto-EM signatures utilized by this invention including the apparatus disclosed and shown above in Figure 1.

25 Using methods known to those skilled in the art, any device capable of detecting and analyzing acoustic energy and/or EM energy through any medium can be used to detect the resonant acoustic and/or acousto-EM signatures utilized by the invention such as disclosed and shown above in Figure 2.

The system shown in Figure 12 gives a schematic overview of the necessary 30 components to be utilized in determining resonant acoustic frequencies of different inorganic materials or structures. Predetermination of the specific frequencies and acoustic and/or

That which is claimed is:

- 5 1. A method for targeting a biologic structure to augment its function which comprises irradiating the biologic structure with acoustic energy having a frequency near or at the resonant frequency of the biologic structure to induce acoustic resonance therein.
2. The method according to claim 1, further comprising determining an acoustic signature of the biologic structure.
- 10 3. A method for targeting a biologic structure to affect at least one function which comprises irradiating the biologic structure with acoustic energy having a frequency near or at the resonant frequency of the biologic structure to induce acoustic resonance therein, and determining at least one acoustic signature and at least one acousto-EM signature of the biologic.
- 15 4. The method according to claim 1, further comprising irradiating the biologic structure with at least one acousto-EM signature of the biologic structure.
5. The method according to claim 1, wherein said acoustic energy is applied at a sufficient power intensity to augment at least one function of the biologic structure, said at least one function being selected from the group of functions consisting of growth, reproduction,
20 regeneration, embryogenesis, metabolism, fermentation, germination, oxidation or reduction activity, wound healing and tissue cutting.
6. The method according to claim 3, wherein said at least one function is selected from the group of functions consisting of disruption and augmentation.
7. The method according to claim 6, wherein said disruption comprises at least one
25 function selected from the group of functions consisting of (a) structural failure of at least

one component in the biologic structure, (b) inhibition of vital processes required for growth, reproduction, metabolism, virulence, and infectivity; and wherein said augmentation comprises at least one function selected from the group of functions consisting of growth, reproduction, regeneration, embryogenesis, metabolism, fermentation, germination, oxidation or reduction activity, wound healing and tissue cutting and (c) lysis, shattering, rupture and inactivation.

8. A method for targeting a specific biologic structure to affect at least one function of the biologic structure comprising:

irradiating the biologic structure with at least one electromagnetic (EM) property and/or field to result in acoustic energy having a frequency including said at least one resonant acoustic frequency of the biologic structure, the acoustic energy being present in an amount sufficient to affect at least one function of the biologic structure.

9. The method according to claim 8, wherein said at least one function is selected from the group of functions consisting of disruption and augmentation.

10. The method according to claim 8, wherein said biologic structure comprises at least one structure selected from the group of structures consisting of virus, bacteria, fungi, tissue masses, worms, arthropods, chitins, plants, animals, microorganisms, multicellular organisms, protozoa, liver, muscle, feet, brain, kidney, spleen, blood, lung, lens of eye, aqueous humor, vitreous humor, animal cell, plant cell, proteins, molecules, cell wall, capsule, spore, pili, plasma membrane, organ, portions of structures, components of structures flagellum, cytoplasmic inclusion body, basal body, parasite, appendages, skin, shell, egg, cement/cement plate and bone.

1.1. A method for specifically targeting a biologic structure and affecting at least one function of the biologic structure by inducing acoustic resonance therein comprising:

a) applying at least two energies selected from the group consisting of at least one acoustic energy and at least one electromagnetic energy, wherein at least one of said at least two energies result in said biologic structure being in acoustic resonance and at least a second

of said at least two energies provides additional energy to said biologic structure; and

b) applying said at least two energies such that a power intensity level is achieved to induce acoustic resonance within the targeted biologic structure and to affect at least one function therein.

5 12. The method according to claim 11, wherein said at least one function comprises at least one function selected from the group of functions consisting of augmenting and disrupting.

13. The method according to claim 11, wherein each of said at least two energies results in acoustic resonance within the targeted biologic.

10 14. The method according to claim 11, wherein said at least two energies comprise at least two energies selected from the group consisting of direct current, alternating current, electric field, magnetic field, electromagnetic radiation and acoustic energy.

15. The method according to claim 14, wherein a frequency of the alternating current is applied to the structure.

15 16. A method for targeting a biologic structure to affect at least one function of the biologic structure comprising applying electromagnetic energy to the biologic structure to induce acoustic resonance therein and affect said at least one function.

17. The method according to claim 16, wherein said electromagnetic energy comprises at least one source selected from the group consisting of at least one electromagnetic energy pattern of the biologic structure energy equivalent in frequency to at least one resonant
20 acoustic frequency of the structure, at least one acousto-EM signature and at least one resonant acousto-EM energy.

18. The method according to claim 16, wherein said electromagnetic energy is applied at a power output level sufficient to affect at least one function of the biologic structure, said
25 at least one function being selected from the group consisting of augmentation and disruption.

19. The method according to claim 16, wherein said electromagnetic energy comprises at least one energy selected from the group consisting of direct current, alternating current, electric field, magnetic field, electromagnetic radiation, and fields which include waves,
30 current, flux, resistance, potential and radiation.

20. The method according to claim 19, wherein said electromagnetic energy comprises at

least one applied acousto-EM signature of the biologic structure.

21. The method according to claim 16, further comprising determining at least one signature of the biologic selected from the group of signatures consisting of at least one acousto-EM signature and at least one acoustic signature.

5 22. The method according to claim 21, wherein said at least one signature of the biologic is compared to at least one previously determined reference signature.

23. A method to induce acoustic stimulation of a biologic structure to detect and/or identify a biologic structure comprising:

- 10 a) applying to the biologic structure at least one acoustic energy comprising at least one non-resonant frequency to stimulate the biologic structure; and
- b) receiving at least one electromagnetic energy pattern from the structure; and
- c) determining at least one non-resonant electromagnetic signature of the stimulated biologic structure.

15 24. A system for inducing acoustic stimulation of a biologic structure to detect and/or identify a biologic structure comprising:

- a) means for applying to the biologic structure at least one acoustic energy comprising at least one non-resonant frequency to stimulate the biologic structure;
- b) means for receiving at least one electromagnetic energy pattern from the structure; and
- 20 c) means for determining at least one non-resonant electromagnetic signature of the stimulated biologic structure.

25. A method for detecting and/or identifying an inorganic or biologic structure comprising:

- a) inducing acoustic resonance in the structure; and
- 25 b) detecting at least one acousto-EM signature of the structure.

26. The method according to claim 25, further comprising comparing at least one currently determined acousto-EM signature with at least one previously determined acousto-EM signature of the structure.

27. The method according to claim 25, wherein said at least one acousto-EM signature is produced by at least one of acoustic energy and electromagnetic properties and/or fields.

28. The method according to claim 25, wherein acoustic resonance is induced with the

introduction of at least one energy selected from the group consisting of acoustic energy including at least one resonant acoustic frequency of the structure, electromagnetic energy which is substantially equivalent to at least one resonant acoustic frequency of the structure and electromagnetic energy which is substantially equivalent to at least one acousto-EM signature of the structure.

29. A system for identifying a structure by determining at least one resonant acoustic signature of the structure comprising:

- a) means for inducing acoustic resonance in the structure;
- b) means for detecting at least one acoustic signature of the structure; and
- c) means for comparing said at least one acoustic signature of the structure with at least one reference acoustic signature.

30. The system according to claim 29, further comprising a means for detecting at least one acoustic-EM signature of the structure.

31. The system according to claim 30, wherein said structure comprises at least one member selected from the group consisting of inorganic and biologic structures.

32. The system according to claim 29, wherein said means for inducing acoustic resonance in the structure includes at least one signal generating device and at least one transducer.

33. The system according to claim 32, wherein placement of the transducer comprises at least one location selected from the group consisting of on the bottom of a vessel, on the walls of a vessel, in a vessel, intravascularly in the biologic structure, extracorporeally of the biologic structure, in vivo, in vitro, in a hand held probe, a piezoelectric sheet, in a remote control unit and in a scalpel tip.

34. A system for identifying a structure by determining at least one acousto-EM signature of the structure comprising:

- a) means for inducing acoustic resonance in the structure; and
- b) means for detecting said at least one acousto-EM signature of the structure.

35. A system for inducing acoustic resonance in a biologic structure to augment at least one function of the biologic structure comprising:

- a) means for generating at least one acoustic signal;
 - b) means for transmitting said at least one acoustic signal to the biologic structure;
- and

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- c) means for controlling the power level of said at least one acoustic signal to augment at least one function of the biologic structure.
36. A system for inducing acoustic resonance in a biologic structure to affect at least one function of the biologic structure comprising:
- 5 a) means for generating at least one electromagnetic signal; and
- b) means for transmitting said at least one electromagnetic signal to the biologic structure.
37. A system for determining induction of acoustic resonance in a structure comprising:
- a) means for generating electromagnetic energy corresponding to at least one
- 10 acousto-EM signature;
- b) means for transmitting said electromagnetic energy to the structure;
- c) means for receiving at least one signal from the structure when said electromagnetic energy has interacted with the structure; and
- d) means for determining induction of acoustic resonance in the structure.
- 15 38. A method for determining induction of acoustic resonance in a structure comprising:
- a) irradiating the structure with electromagnetic energy corresponding to at least one acousto-EM signature;
- b) receiving at least one signal from the structure when said electromagnetic energy has interacted with the structure; and
- 20 c) determining induction of acoustic resonance in the structure.
39. A method to affect at least one function of a living transducer biologic structure comprising applying at least one electromagnetic energy to the biologic structure, said at least one electromagnetic energy comprising at least one frequency which includes at least one resonant frequency of the biologic structure to induce acoustic resonance within
- 25 the biologic structure, the energy being present in an amount sufficient to affect at least one function of the biologic structure.
40. A method for targeting an inorganic structure to affect said structure, the method comprising applying at least one resonant acousto-EM energy.
41. The method of claim 40, which said structure is affected by disruption.
- 30 42. The method of claim 40, wherein said structure is affected by augmentation.

43. A method for detecting an inorganic structure comprising:
- a) inducing acoustic resonance in the structure; and
 - b) detecting at least one resonant acousto-EM energy.
44. A method for detecting an inorganic structure comprising:
- a) inducing acoustic resonance in the structure by applying an acousto-EM signature; and
 - b) detecting at least one acoustic signature.

INTERNATIONAL SEARCH REPORT

International application No.

PCT/US99/20776

A. CLASSIFICATION OF SUBJECT MATTER

IPC(7) : A61B 17/22, 6/00; A61N 7/00
US CL : 600/427, 429, 439; 601/2, 3, 4; 73/579

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

U.S. : 600/427, 429, 439; 601/2, 3, 4; 607/97, 51; 604/22; 73/579

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)
EAST: resonant, resonance, biologic, biological, cell, acoustic, ultrasonic, ultrasound

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	US 3,774,717 A (CHODOROW) 27 November 1973 (27.11.1973) see abstract, Figures, column 1 line 35 - column 3 line 6	1-2, 23-25, 28, 34
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Y		10, 16-18, 21-22, 35
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A		3-4, 19-20, 27, 36-39
X	US 4,315,514 A (DREWES ET AL.) 16 February 1982 (16.02.1982), see abstract, Figures, column 2 lines 30-64.	1, 5-7, 9-11, 13
---		-----
Y		8, 12, 14-18, 21-22, 35
Y	US 5,413,550 A (CASTEL) 09 May 1995 (09.05.1995) see abstract, Figures, column 1 lines 5-16, column 6 line 55 - column 7 line 27	35
Y	US 5,595,178 A (VOSS ET AL.) 21 January 1997 (21.01.1997) see abstract, Figures, column 1 lines 5-18, column 2 lines 5-29, column 3 line 47 - column 4 line 13, column 4 lines 29-44	8, 10, 12, 14-15
X	US 5,777,228 A (TSUBOI ET AL.) 07 July 1998 (07.07.1998) see abstract, Figures, column 3 line 33 - column 9 line 20	25-26, 28-29, 32-34



Further documents are listed in the continuation of Box C.



See patent family annex.

Special categories of cited documents:	
"A" document defining the general state of the art which is not considered to be of particular relevance	"T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention
"E" earlier application or patent published on or after the international filing date	"X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone
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"O" document referring to an oral disclosure, use, exhibition or other means	"&" document member of the same patent family
"P" document published prior to the international filing date but later than the priority date claimed	

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INTERNATIONAL SEARCH REPORT

International application No.

PCT/US99/20776

C (Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X, P --- A, P	US 5,886,263 A (NATH ET AL.) 23 March 1999 (23.03.1999) see abstract, Figures, column 1 line 33 - column 2 line 17	25-26, 28-29, 32-34 ----- 27, 30-31, 36-38